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Assessing histidine tags for recruiting deoxyribozymes to catalyze peptide and protein modification reactions†

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We evaluate the ability of hexahistidine (His₆) tags on peptide and protein substrates to recruit deoxyribozymes for modifying those substrates. For two different deoxyribozymes, one that creates tyrosine-RNA nucleopeptides and another that phosphorylates tyrosine side chains, we find substantial improvements in yield, k_{obs} , and K_{m} for peptide substrates due to recruiting by His₆/Cu²⁺. However, the recruiting benefits of the histidine tag are not observed for larger protein substrates, likely because the tested deoxyribozymes either cannot access the target peptide segments or cannot function when these segments are presented in a structured protein context.

Introduction

Deoxyribozymes are single-stranded DNA sequences with catalytic activity.^{1–5} Our laboratory seeks deoxyribozymes for covalent protein modification.⁶ In these efforts, a key challenge is achieving DNA-catalyzed reactions using relatively low protein concentrations that enable preparative utility. We recently described the use of azide-functionalized peptide substrates during *in vitro* selection, which enabled identification of deoxyribozymes with peptide K_{m} of ~100 μM .⁷ However, that approach is not applicable to all DNA-catalyzed reactions. Here we pursued an alternative and potentially more general strategy: recruiting deoxyribozymes for modifying peptides or proteins using histidine tags.

Rosen *et al.* recently reported that metal ion-mediated interactions between a suitably functionalized DNA oligonucleotide and a metal-binding (*e.g.*, histidine-tagged) protein can be used to recruit the DNA oligonucleotide to a specific region on a protein surface (Fig. 1A).⁸ Subsequent nonenzymatic, DNA-templated reaction leads to site-specific protein modification by attachment of a DNA strand, whereas many other chemical modification approaches for proteins are not site-selective.^{9–23} Here, we evaluated experimentally whether a histidine tag can be used to recruit a deoxyribozyme—which in this context is simply a long DNA oligonucleotide—to a peptide or protein, thereby enhancing DNA-catalyzed covalent modification of

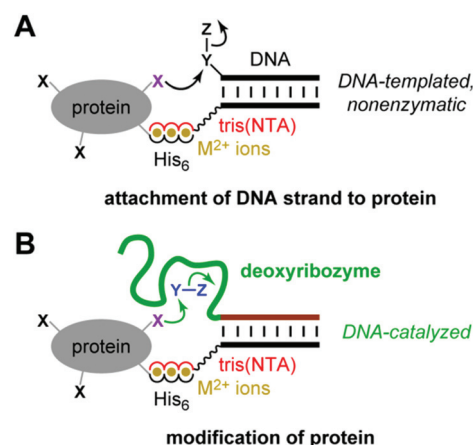


Fig. 1 Use of histidine tags to recruit reagents or catalysts for peptide and protein modification. (A) The reported use of hexahistidine (His₆) tags and divalent metal ions (M²⁺) for directing nonenzymatic, DNA-templated reaction of protein side chains.⁸ In one case, the authors instead used a natively metal-binding protein. NTA = nitrilotriacetic acid. (B) As evaluated in this study, the potential use of His₆ tags and M²⁺ to recruit a deoxyribozyme for catalyzing peptide and protein modification. See Fig. 2 for details of DNA catalysis arrangement, including reaction partner for the protein (shown here as Y–Z), as well as NTA structure.

that substrate, in principle by any modification reaction that can be catalyzed by DNA (Fig. 1B). Unlike the nonenzymatic, strictly proximity-driven approach of Fig. 1A, the strategy of Fig. 1B uses Watson-Crick base pairs for recruiting the deoxyribozyme to the peptide or protein substrate. The subsequent modification reaction still requires enzymatic catalysis by the DNA.

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Results

Evaluating the His₆-tag recruiting strategy with the 8XJ105 deoxyribozyme, which forms peptide-RNA conjugates

For evaluating the histidine tag recruiting strategy of Fig. 1B, we initially used the 8XJ105 deoxyribozyme, which attaches a 5'-triphosphorylated RNA oligonucleotide (pppRNA) to a tyrosine hydroxyl group in a peptide substrate, forming a nucleopeptide (peptide-RNA) conjugate.⁷ Such peptide-RNA (or peptide-DNA) conjugates are integrally involved in many key biological processes,^{24–32} and they are useful for various applications as well.^{33–40} 8XJ105 was identified by an *in vitro* selection strategy that uses an untethered (discrete, free) peptide during each selection round. As a consequence, 8XJ105 has substantial activity with an untethered peptide, but only at very high peptide concentration (apparent peptide $K_m > 1$ mM). 8XJ105 tolerates a wide variety of sequence contexts for the substrate's tyrosine residue, *i.e.*, 8XJ105 is peptide sequence-general. Therefore, we sought to use histidine tags to recruit 8XJ105 for conjugating RNA to a tyrosine-containing peptide.

A DNA anchor oligonucleotide was synthesized, bearing at either its 3'-end or 5'-end a tris(NTA) moiety that efficiently binds certain divalent metal ions (NTA = nitrilotriacetic acid; Fig. 2A). As was done by Rosen *et al.*,⁸ we adapted the synthetic procedure from the work of Goodman *et al.*⁴¹ The deoxyribozyme interacts with the DNA anchor by standard Watson-Crick base pairs, such that recruiting the DNA anchor to the peptide substrate also recruits the deoxyribozyme. The peptide sub-

strate bears at its N-terminus a hexahistidine (His₆) tag. As shown in Fig. 2B, the intention is that the tris(NTA) and His₆ moieties bind simultaneously to divalent metal ions (M^{2+}), thereby recruiting the deoxyribozyme to the peptide and enabling DNA-catalyzed peptide modification.

The 8XJ105 deoxyribozyme was first evaluated for its catalytic activity in control experiments with His₆-tagged peptide substrates in the absence of His₆/ M^{2+} recruiting (Fig. 3A, controls). These experiments are important to establish the conditions and substrates with which the recruiting effect will then be evaluated. 8XJ105 requires 1 mM Zn^{2+} as a cofactor (as well as 40 mM Mg^{2+} and 20 mM Mn^{2+}). Therefore, the controls used 1 mM Zn^{2+} and a DNA anchor oligonucleotide that lacks the tris(NTA) modification. 8XJ105 showed very little activity with the 11-mer peptide H₆AAYAA (2% yield in 16 h at 10 or 30 μ M peptide and <0.5% yield at 100 μ M peptide). Zn^{2+} leads to peptide aggregation upon interaction with the His₆ tag, which constitutes more than half of the peptide. Such aggregation was visible by eye as a white precipitate at high peptide concentration (1 mM). 8XJ105 was more active with two longer 18-mer and 24-mer His₆-tagged peptides, each containing a single tyrosine residue. The 24-mer led to 11% yield in 16 h at 100 μ M peptide, although reduced activity was still observed at even higher peptide concentrations (>100 μ M). The 11% yield at 100 μ M can be compared with 5% yield observed with a non-His₆-tagged 24-mer peptide at the same concentration. The 24-mer peptide was therefore used as the substrate to assess the particular recruiting strategy of Fig. 2B, which is expected to reduce the required peptide concentration and increase the reaction yield of peptide-RNA conjugation.

The third-row metal ions Co^{2+} , Ni^{2+} , Cu^{2+} , and Zn^{2+} interact efficiently with NTA and have been used in Immobilized-Metal Affinity Chromatography (IMAC) for applications such as protein purification.^{42–45} We tested each of these metal ions with 8XJ105 according to the recruiting strategy of Fig. 2B. The 3'-tris(NTA)-modified DNA anchor oligonucleotide was incubated with 8XJ105, pppRNA, the His₆-tagged 24-mer peptide substrate, and 10 μ M of one of Co^{2+} , Ni^{2+} , or Cu^{2+} , noting again that 1 mM Zn^{2+} is present in all experiments [Fig. 3A, 3'-tris(NTA) anchor data]. Inclusion of the tris(NTA) moiety but no extra M^{2+} led to 2.1-fold increase in yield (from 11% to 23%) relative to the non-tris(NTA) control at 100 μ M peptide. No additional benefit arose by including 10 μ M Co^{2+} or Ni^{2+} . In contrast, including 10 μ M Cu^{2+} led to 6.4-fold increase in yield (from 6.9% to 44%) relative to control at 30 μ M peptide. A yield of 31% was observed for a non-His₆-tagged 24-mer peptide, albeit only at 30-fold higher peptide concentration (>1 mM; yields decreased substantially at high peptide concentration, again attributed to aggregation). Therefore, a histidine tag successfully recruits the 8XJ105 deoxyribozyme to modify its peptide substrate. Among the evaluated divalent metal ions, Cu^{2+} was most effective, consistent with its tightest binding in metal complexes.⁴⁶ The alternative 5'-tris(NTA) on the anchor oligonucleotide led to 2.8-fold yield increase relative to non-tris(NTA) control at 100 μ M peptide, but none of Co^{2+} , Ni^{2+} , or Cu^{2+} provided any additional yield increase

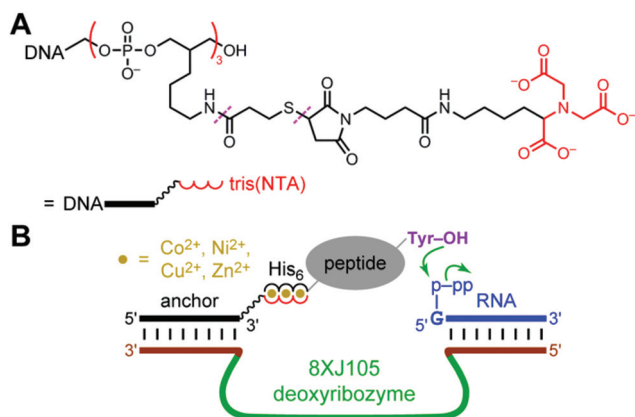


Fig. 2 Strategy for using histidine tags to recruit a deoxyribozyme to a peptide substrate and enable DNA-catalyzed peptide modification. (A) Tris(NTA)-modified DNA anchor oligonucleotide. The 3'-tris(NTA) moiety was connected to the DNA by a hexa(ethylene glycol) spacer. For the 5'-tris(NTA)-modified DNA anchor, the 5'-tris(NTA) moiety was connected to the DNA by a T₁₅ oligonucleotide spacer. Each tris(NTA)-modified oligonucleotide was synthesized by a multistep procedure in which the linkages marked with dashes were created (see the Experimental section). (B) Design showing the intended interactions among the 8XJ105 deoxyribozyme that forms nucleopeptide linkages, 3'-tris(NTA)-modified DNA anchor, His₆-tagged peptide substrate, and divalent metal ions. Success is revealed by either or both of increase in peptide-RNA conjugation yield and leftward shift in peptide concentration dependence.



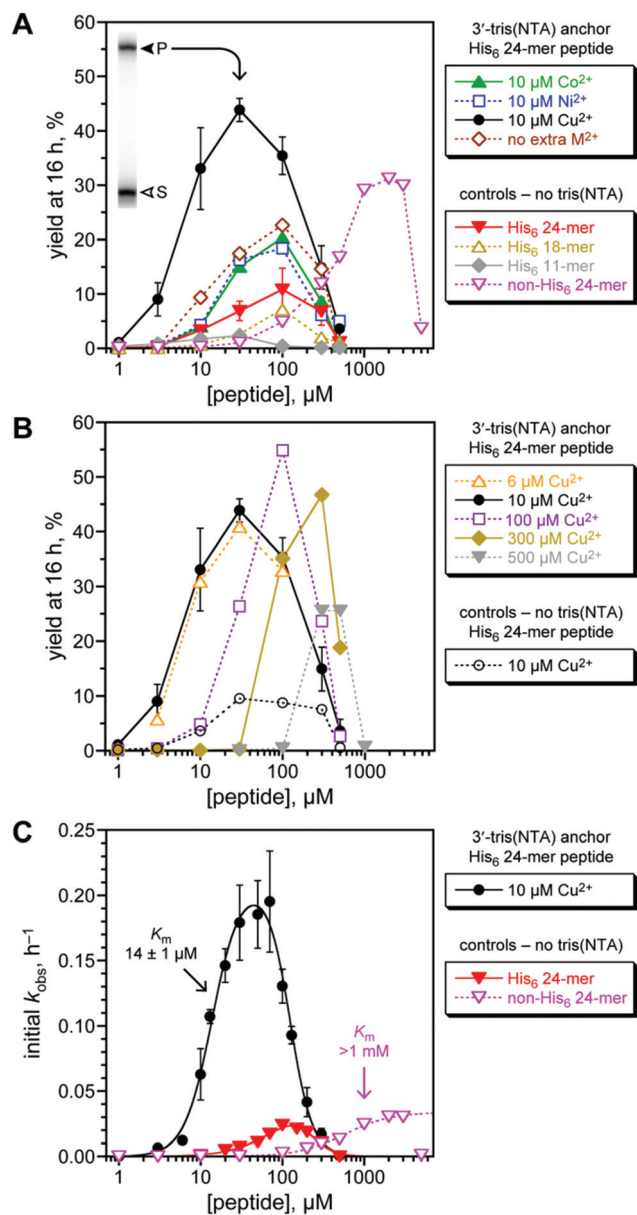


Fig. 3 Assessing histidine tag recruiting for the 8XJ105 deoxyribozyme. See the Experimental section for explanation of error bars in all panels. Conditions: 20 nM 3'-³²P-radiolabeled 5'-pppRNA, 0.5 μM 8XJ105 deoxyribozyme, 2 μM 3'-tris(NTA) DNA anchor oligonucleotide, 1–5000 μM peptide, 70 mM HEPES, pH 7.5, 40 mM MgCl₂, 20 mM MnCl₂, 1 mM ZnCl₂, 150 mM NaCl, and CoCl₂, NiCl₂, or Cu(NO₃)₂ as indicated at room temperature. CuCl₂ gave equivalent outcome. (A) Yield at 16 h of peptide-RNA conjugate as a function of concentration of His₆-tagged 24-mer peptide (or untagged 24-mer in one case). All experiments including the controls were performed with 1 mM Zn²⁺; 10 μM Co²⁺, Ni²⁺, or Cu²⁺ was additionally present where indicated. *Inset*: PAGE data at 16 h for 3'-³²P-radiolabeled RNA and 30 μM His₆-tagged 24-mer peptide with Cu²⁺ (S = substrate, P = product). See Fig. S1 (ESI[†]) for data with 5'-tris(NTA) moiety. 24-mer H₆SAGERASAEDMARAAYAA (for non-His₆, replace H₆ with ASAASA); 18-mer H₆SAEDMARAAYAA; 11-mer H₆AAYAA. (B) Optimization of Cu²⁺ concentration, and demonstration that Cu²⁺ alone [without tris(NTA)] is not responsible for the recruiting effect. (C) Determination of peptide *K*_m values using initial-rate kinetics. See curve fit descriptions and fit details in the Experimental section.

(Fig. S1, ESI[†]), indicating a smaller recruiting benefit from this 5'-tris(NTA) alternative.

The Cu²⁺ concentration was optimal at 10 μM when the 3'-tris(NTA) anchor oligonucleotide was at 2 μM, such that the metal-binding capacity is 6 μM (Fig. 3B). As an important control experiment, without the 3'-tris(NTA) moiety, 10 μM Cu²⁺ supported catalysis with no greater efficiency than in the absence of Cu²⁺.

The substantial recruiting effect of His₆/Cu²⁺ was confirmed by evaluating the rate of DNA-catalyzed peptide modification (Fig. 3C). From initial-rate kinetics, *k*_{obs} values for the His₆-tagged 24-mer peptide substrate were obtained both with and without 3'-tris(NTA) on the anchor oligonucleotide. Although generic inhibition of activity at >100 μM peptide was observed, the data again reveal a strong recruiting effect, with peptide *K*_m of 14 μM. Comparing the data for the His₆-tagged 24-mer and untagged non-His₆ 24-mer peptides indicates >70-fold decrease in *K*_m value due to recruiting.

Evaluating the His₆-tag recruiting strategy with the 6CF134 tyrosine kinase deoxyribozyme

In separate experiments, we investigated the Fig. 1B recruiting strategy for DNA-catalyzed tyrosine side chain phosphorylation (kinase activity). Tyrosine phosphorylation is one of many important, natural protein post-translational modification reactions,^{47–52} and synthetic tyrosine kinase enzymes will have substantial utility. We previously reported the 6CF134 tyrosine kinase deoxyribozyme, which phosphorylates a short hexapeptide substrate when that substrate is covalently tethered to a DNA anchor oligonucleotide (Fig. 4A).⁵³ However, 6CF134 does not catalyze detectable phosphorylation of a discrete, untethered peptide at any peptide concentration tested. Here we sought to recruit 6CF134 to an untethered 24-mer peptide substrate *via* a His₆ tag. In the presence of 100 μM Cu²⁺, 88% conversion of phosphorylated peptide was observed by MALDI mass spectrometry when the 3'-tris(NTA) was included (Fig. 4B), compared to 50% yield for 6CF134 with a tethered peptide under similar conditions (Fig. 4A). Little or no phosphorylation was observed in negative control assays that omitted one of 3'-tris(NTA) (0.9% conversion; Fig. 4B), Cu²⁺ (0.3%; data not shown), or the pppRNA phosphoryl donor (<0.05%; data not shown), or when the non-His₆ 24-mer peptide was used (0.09%; data not shown). From these observations, we conclude that 6CF134 is inherently capable of modifying an untethered peptide, despite its strict tether requirement found in our original report.⁵³ The His₆ tag and tris(NTA) interaction functionally replaces the covalent tether and enables 6CF134 catalysis with the formally untethered peptide substrate.

Evaluating the His₆-tag recruiting strategy with protein substrates

Rosen *et al.* used both His₆-tagged and natively metal-binding proteins in their work.⁸ We evaluated the recruiting strategy of Fig. 2B for 8XJ105, using two His₆-tagged proteins (lysozyme and plasminogen activator inhibitor 1 [PAI-1]) and one natively



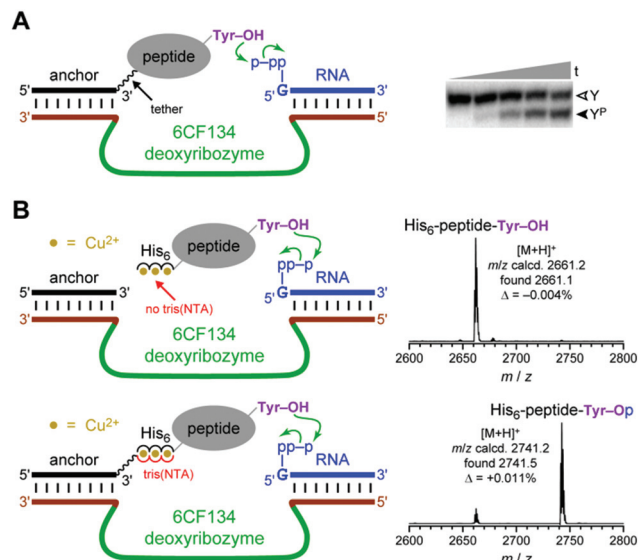


Fig. 4 The 6CF134 tyrosine kinase deoxyribozyme and its His₆ recruiting to a peptide substrate. (A) Originally reported arrangement of 6CF134 and disulfide-tethered peptide substrate. No phosphorylation is observed without the tether;⁵³ ~50% phosphorylation yield is observed with the tether. Conditions: 20 nM 5'-³²P-radiolabeled DNA-anchored CAAYAA peptide, 0.5 μM 6CF134 deoxyribozyme, 5 μM 5'-pppRNA, 70 mM HEPES, pH 7.5, 40 mM MgCl₂, 20 mM MnCl₂, 0.5 mM ZnCl₂, and 150 mM NaCl at room temperature (*t* = 30 s, 30 min, 2 h, 5 h, and 16 h; Y = substrate, Y^P = product). (B) MALDI mass spectrometry reveals successful recruiting of 6CF134 to phosphorylate the 24-mer His₆-tagged peptide substrate (same substrate as in Fig. 3). Conditions: 30 μM peptide, 30 μM DNA anchor oligonucleotide with or without 3'-tris(NTA), 35 μM 6CF134 deoxyribozyme, 40 μM 5'-pppRNA, 70 mM HEPES, pH 7.5, 40 mM MgCl₂, 20 mM MnCl₂, 2 mM ZnCl₂, 150 mM NaCl, and 100 μM Cu(NO₃)₂ at room temperature for 20 h.

metal-binding protein (carboxypeptidase B). For all three proteins and using 3'-³²P-radiolabeled pppRNA, no modification of any tyrosine residues by 8XJ105 was observed by PAGE-shift assay (Fig. S2, ESI†). Separately, we also sought to use 6CF134 with lysozyme and PAI-1, albeit unsuccessfully, as assayed by trypsin digestion and MALDI mass spectrometry (Fig. S3, ESI†).

Discussion

In this study, we performed experiments to evaluate the histidine tag recruiting strategy of Fig. 1B for enhancing DNA-catalyzed peptide and protein modification. Substantially higher peptide modification yields and *k*_{obs} values were observed, along with substantial effects on peptide *K*_m values. The His₆/Cu²⁺ recruiting effect allowed, for the first time, successful DNA-catalyzed tyrosine phosphorylation of a discrete, untethered peptide substrate. However, where tested, DNA-catalyzed protein modification using the recruiting strategy was not observed. Unlike nonenzymatic protein modification reactions such as those studied by Rosen *et al.*,⁸ where simple proximity is responsible for the enhanced reactivity (Fig. 1A), achieving

DNA-catalyzed reactions of proteins faces additional challenges, even with the benefit of histidine tag recruiting. For example, access by a deoxyribozyme to residues on a structured protein's surface may be restricted, and the recruiting effect may not overcome this problem. In addition, a deoxyribozyme may not function well (or at all) with peptide segments on a protein's surface, if those segments are unable to adopt a conformation suitable to allow their DNA-catalyzed modification.

For deoxyribozymes that do function well with protein substrates, the recruiting strategy should allow site-selective modification of particular surface-exposed residues, based on the geometric relationship among the tris(NTA) recruiting site, the recruited deoxyribozyme, and the side chains themselves (see Fig. 1B). This directed reactivity strategy for a sequence-general deoxyribozyme differs from our recently reported experiments in which inherently sequence-selective deoxyribozymes were identified using particular peptide sequences during the selection process, such that the resulting deoxyribozymes function only with those peptide sequences.^{7,54} We are currently seeking protein-modifying deoxyribozymes using various approaches, and with such DNA catalysts we intend to evaluate the histidine tag recruiting strategy to achieve site selectivity. The experiments reported here provide an encouraging framework for these future studies.

Experimental

Oligonucleotides and oligonucleotide-tris(NTA) conjugates

General considerations. DNA oligonucleotides were obtained from Integrated DNA Technologies (Coralville, IA) or prepared by solid-phase synthesis on an ABI 394 instrument using reagents from Glen Research. 5'-Triphosphorylated RNA (pppRNA) oligonucleotides were prepared by *in vitro* transcription using synthetic DNA templates and T7 RNA polymerase.⁵⁵ All oligonucleotides were purified by 7 M urea denaturing PAGE with running buffer 1× TBE (89 mM each Tris and boric acid and 2 mM EDTA, pH 8.3) as described previously.^{56,57} Sequences are collected in Table S1 (ESI†).

Procedure for synthesis of tris(NTA)-modified DNA anchor. DNA oligonucleotides with three primary amino groups at the 3'-end or 5'-end were synthesized using the Uni-Link Amino Modifier (Clontech). The procedure for incorporating tris(NTA) was adapted from Goodman *et al.*⁴¹ A 50 μL sample containing 10 nmol of tris(amino)-modified DNA oligonucleotide in 100 mM sodium phosphate buffer, pH 7.5, 100 mM NaCl, and 25 mM SPDP [*N*-succinimidyl 3-(2-pyridylthio)propionate, Fisher cat. no. PI21857] was incubated at room temperature for 1 h. The SPDP was added from a freshly prepared 100 mM stock in DMF. A desalting column (Micro Bio-Spin P-6, Bio-Rad) was washed with 100 mM sodium phosphate buffer, pH 7.5, and used to remove excess SPDP from the sample. The eluted sample was treated with 6.3 μL of 100 mM TCEP [tris(carboxyethyl)phosphine hydrochloride, Chem-Impex] and incubated at room temperature for 30 min. Excess TCEP was



removed with another desalting column. The eluted sample was treated with 7 μL of 100 mM maleimido- C_3 -NTA (Dojindo cat. no. M035-10), 1 μL of 1 M sodium phosphate buffer, pH 7.5, and 2.5 μL of 3 M NaCl, and the sample was incubated at room temperature for 1 h. The sample was purified by 20% PAGE. The product was extracted from the gel in 10 mM Tris, pH 8.0 and 300 mM NaCl (no EDTA) and precipitated with ethanol. Typical yield after gel extraction and precipitation was 20–30%. The product was analyzed by mass spectrometry on a Bruker UltrafleXtreme MALDI-TOF mass spectrometer with matrix 3-hydroxypicolinic acid in positive ion mode at the UIUC School of Chemical Sciences Mass Spectrometry Laboratory. For the 3'-tris(NTA) DNA anchor oligonucleotide used in peptide-RNA conjugation, $[\text{M} + \text{H}]^+$ calcd 7341.6, found 7340.6 ($\Delta = -0.014\%$). For the 5'-tris(NTA) DNA anchor oligonucleotide used in peptide-RNA conjugation, $[\text{M} + \text{H}]^+$ calcd 11 560.3, found 11 564.6 ($\Delta = +0.037\%$). For the 3'-tris(NTA) DNA anchor oligonucleotide used in peptide tyrosine phosphorylation, $[\text{M} + \text{H}]^+$ calcd 7343.7, found 7343.3 ($\Delta = -0.005\%$). Images of the mass spectra are shown in Fig. S4 (ESI[†]).

Synthesis of peptides

Peptides were prepared by solid-phase synthesis using Fmoc Rink amide MBHA resin on a Liberty Automated Microwave Peptide Synthesizer (CEM Corporation) using the instrument's standard wash procedures. Each synthesis was performed at 0.1 mmol scale, initiated using 130 mg of Rink amide resin with a loading capacity of 0.77 mmol g^{-1} (Chem-Impex). For each Fmoc deprotection step, 20% piperidine in DMF with 0.1 M HOBt was used at 30 W of microwave power and 75 $^{\circ}\text{C}$ for 3 min. Each coupling used 5 equivalents (2.5 mL of 0.2 M in DMF, 0.5 mmol) of Fmoc-amino acid, 5 equivalents (1 mL of 0.5 M in DMF, 0.5 mmol) of HCTU (*N,N,N',N'*-tetramethyl-*O*-(1*H*-6-chlorobenzotriazol-1-yl)uronium hexafluorophosphate), and 10 equivalents (0.5 mL of 2 M in *N*-methylpyrrolidone, 1 mmol) of DIPEA (*N,N*-diisopropylethylamine). All amino acid monomers except Arg and His were coupled twice at 20 W and 75 $^{\circ}\text{C}$ for 5 min. For Arg, to suppress δ -lactam formation, the first coupling was performed without microwaver power at room temperature for 25 min, followed by 25 W at 75 $^{\circ}\text{C}$ for 5 min; the second coupling was performed at 20 W and 75 $^{\circ}\text{C}$ for 5 min. For His, to suppress racemization, both couplings were performed without microwave power at room temperature for 2 min followed by 25 W at 50 $^{\circ}\text{C}$ for 4 min. After each coupling step as well as the final Fmoc deprotection step, capping was performed using 7 mL of 0.5 M acetic anhydride and 0.125 M DIPEA in DMF at 40 W and 65 $^{\circ}\text{C}$ for 2 min. The peptide was cleaved from the solid support by stirring the resin in a separate vial with a solution containing 5 mL of tri-fluoroacetic acid (TFA), 125 μL of water, and 50 μL of triisopropylsilane for 90 min. The liquid solution was separated from the resin by filtration. This solution was dried on a rotary evaporator, providing an oily solid. To this material was added 20 mL of cold diethyl ether, and the peptide was obtained as a white solid that was filtered and purified by HPLC. Each

peptide had an N-terminal acetyl group and C-terminal amide group.

His₆ 11-mer: HHHHHHAAYAA

His₆ 18-mer: HHHHHHSAEDMARAYAA

His₆ 24-mer: HHHHHHSAGERASAEDMARAYAA

non-His₆ 24-mer: ASAASASAGERASAEDMARAYAA

Single-turnover deoxyribozyme assay and mass spectrometry procedures

General considerations. Metal ions MgCl_2 , MnCl_2 , CoCl_2 , NiCl_2 , and $\text{Cu}(\text{NO}_3)_2$ were added from 10 $\times$ stock solutions, which were diluted from 1 M (or 500 mM for NiCl_2) solutions. ZnCl_2 was added from a 10 $\times$ stock solution containing 10 mM ZnCl_2 , 20 mM HNO_3 , and 200 mM HEPES at pH 7.5; this stock was freshly prepared from a solution of 100 mM ZnCl_2 in 200 mM HNO_3 . 3'-³²P-radiolabeling of 5'-pppRNA was achieved starting from the unlabeled 5'-pppRNA using 5'-³²P-pCp and T4 RNA ligase.⁵⁸

Peptide-RNA conjugation by the 8XJ105 deoxyribozyme. A 5 μL sample containing 20 pmol of DNA anchor oligonucleotide [with or without 3'-tris(NTA) as appropriate], one of 100 pmol of Co^{2+} , 100 pmol of Ni^{2+} , or 60–5000 pmol of Cu^{2+} if applicable, and peptide substrate was incubated in 50 mM HEPES, pH 7.5, and 150 mM NaCl at room temperature for 1 h (this preincubation step was later determined to be dispensable). The reaction was initiated by bringing the sample to 10 μL total volume containing 20 nM 3'-³²P-radiolabeled 5'-pppRNA, 0.5 μM 8XJ105 deoxyribozyme, 2 μM DNA anchor oligonucleotide, 1–5000 μM peptide, 70 mM HEPES, pH 7.5, 40 mM MgCl_2 , 20 mM MnCl_2 , 1 mM ZnCl_2 , and 150 mM NaCl. For Co^{2+} or Ni^{2+} as the recruiting metal ion, the final concentration was 10 μM ; for Cu^{2+} , 6–500 μM . The sample was incubated at room temperature, and 2 μL aliquots were quenched at appropriate times with 5 μL of stop solution (80% formamide, 1 $\times$ TBE [89 mM each Tris and boric acid, 2 mM EDTA, pH 8.3], 50 mM EDTA, and 0.025% each bromophenol blue and xylene cyanol). To each aliquot before PAGE was added 10 pmol of a decoy oligonucleotide (complementary to the deoxyribozyme's 40 nt catalytic region along with 10 nt of binding arms on either side), to displace the deoxyribozyme from the substrate and product. Samples were separated by 20% PAGE and quantified with a PhosphorImager.

Peptide tyrosine phosphorylation by the 6CF134 deoxyribozyme. The assay of Fig. 4A was performed as described.⁵³ The assays of Fig. 4B were performed with much higher concentrations of DNA anchor oligonucleotide, unirradiolabeled 5'-pppRNA, and recruiting metal ion, to enable peptide product characterization by mass spectrometry. A 5 μL sample containing 300 pmol of DNA anchor oligonucleotide [with or without 3'-tris(NTA) as appropriate], 1 nmol of $\text{Cu}(\text{NO}_3)_2$, and 300 pmol of His₆-tagged 24-mer peptide substrate was incubated in 50 mM HEPES, pH 7.5, and 150 mM NaCl at room temperature for 1 h (this preincubation step was later determined to be dispensable). The reaction was initiated by bringing the sample to 10 μL total volume containing 30 μM His₆-tagged 24-mer peptide, 30 μM DNA anchor oligonucleotide, 35 μM 6CF134



deoxyribozyme, 40 μM uniradiolabeled 5'-pppRNA, 70 mM HEPES, pH 7.5, 40 mM MgCl_2 , 20 mM MnCl_2 , 2 mM ZnCl_2 , 150 mM NaCl, and 100 μM $\text{Cu}(\text{NO}_3)_2$. The sample was incubated at room temperature for 20 h, immediately desalted by Millipore C₁₈ ZipTip, and analyzed by MALDI mass spectrometry. Data were acquired on a Bruker UltrafleXtreme MALDI-TOF mass spectrometer with matrix 2,5-dihydroxybenzoic acid in positive ion mode.

Analysis of deoxyribozyme assay data

Data for the control experiment with the His₆ 24-mer (Fig. 3A and Fig. S1, ESI†) are shown as mean $\pm$ SD ($n = 4$). Data for the experiment with the 3'-tris(NTA) DNA anchor oligonucleotide with 10 μM Cu^{2+} (Fig. 3A and B) are shown as mean $\pm$ SD ($n = 3$ –5). Data for the initial-rate kinetics with 10 μM Cu^{2+} and the control experiment with the His₆ 24-mer in Fig. 3C are shown as mean $\pm$ SD ($n = 3$); in the latter case, the error bars are smaller than the data points. All other data are $n = 1$.

Initial-rate kinetics data for Fig. 3C were obtained by fitting the initial linear portion of plots (yield *versus* time up to 2 h) to a straight line. The peptide K_m values were determined by fitting the data to $k_{\text{obs}} = k_{\text{max}} \cdot [C^n / (K_m^n + C^n)] \cdot [1 - C^m / (K_i^m + C^m)]$, where C is the peptide concentration, k_{max} is the k_{obs} at saturating peptide concentration, K_m is the K_m value for productive peptide binding, K_i is the inhibition constant for unproductive peptide binding, and n and m are Hill coefficients for productive and unproductive peptide binding, respectively. The K_m value for the non-His₆ 24-mer was determined by fitting the data to $k_{\text{obs}} = k_{\text{max}} \cdot [C^n / (K_m^n + C^n)]$. Curve fit values were as follows. For the 3'-tris(NTA) anchor with Cu^{2+} , $K_m = 14 \pm 1 \mu\text{M}$, $K_i = 124 \pm 8 \mu\text{M}$, $n = 2.6 \pm 0.4$ and $m = 3.2 \pm 0.5$. For the control experiment with the His₆ 24-mer, $K_m = 82 \pm 75 \mu\text{M}$, $K_i = 218 \pm 68 \mu\text{M}$, $n = 1.6 \pm 0.6$ and $m = 3.3 \pm 1.0$ (these values are not necessarily interpretable due to the large errors; K_m is likely to be $\gg 80 \mu\text{M}$). For the control using non-His₆ 24-mer, $K_m = 579 \pm 57 \mu\text{M}$ and $n = 1.5 \pm 0.2$ (the 5 mM data point was omitted from the fit; K_m is likely to be $> 1 \text{ mM}$).

Assays of 8XJ105 with protein substrates

Following the approach of Rosen *et al.*,⁸ we assayed the 8XJ105 deoxyribozyme with each of three different proteins as potential substrates. His₆-lysozyme was from US Biological (cat. no. 155733, 15.2 kDa). His₆-plasminogen activator inhibitor 1 (PAI-1) was from US Biological (cat. no. 170035, 45 kDa). Carboxypeptidase B, which natively binds metal ions, was from Worthington Biochemical Corporation (cat. no. LS005305, 34.5 kDa). A 5 μL sample containing 20 pmol of 3'- or 5'-tris(NTA) DNA anchor oligonucleotide, 100–3000 pmol of $\text{Cu}(\text{NO}_3)_2$, and 100–1000 pmol of protein substrate was incubated in 50 mM HEPES, pH 7.5, 150 mM NaCl, and 0.04% (v/v) Tween-20 at room temperature for 1 h. The reaction was initiated by bringing the sample to 10 μL total volume containing 20 nM 3'-³²P-radiolabeled 5'-pppRNA, 0.5 μM 8XJ105 deoxyribozyme, 2 μM 3'- or 5'-tris(NTA) DNA anchor oligonucleotide, 10–100 μM protein, 70 mM HEPES, pH 7.5, 1 or 40 mM MgCl_2 , 5 or 20 mM MnCl_2 , 0.5 or 1 mM ZnCl_2 ,

150 mM NaCl, 10–300 μM $\text{Cu}(\text{NO}_3)_2$, and 0.02% (v/v) Tween-20. The sample was incubated at room temperature for 16 h. In all three cases, 16% SDS-PAGE revealed no detectable product formation ($< 0.5\%$; Fig. S2, ESI†).

Assays of 6CF134 with protein substrates

We assayed the 6CF134 deoxyribozyme with each of lysozyme and PAI-1 as potential substrates. A 5 μL sample containing 150 or 300 pmol of 3'-tris(NTA) DNA anchor oligonucleotide, 500 or 1000 pmol of $\text{Cu}(\text{NO}_3)_2$, and 100 or 300 pmol of protein substrate was incubated in 50 mM HEPES, pH 7.5 and 150 mM NaCl at room temperature for 1 h. The reaction was initiated by bringing the sample to 10 μL total volume containing 10 or 30 μM protein, 15 or 30 μM 3'-tris(NTA) DNA anchor oligonucleotide, 20 or 35 μM 6CF134 deoxyribozyme, 25 or 40 μM uniradiolabeled 5'-pppRNA, 70 mM HEPES, pH 7.5, 40 mM MgCl_2 , 0 or 20 mM MnCl_2 , 0.5 or 2 mM ZnCl_2 , 150 mM NaCl, and 50 or 100 μM $\text{Cu}(\text{NO}_3)_2$. The sample was incubated at room temperature for 20 or 48 h. Trypsin digestion was performed to prepare the sample for mass spectrometry. The sample was reduced and denatured by adding 10 μL of 6 M urea in 100 mM Tris-HCl, pH 8.0, and 1 μL of 250 mM DTT and incubating the sample at 37 $^\circ\text{C}$ for 1 h. To block cysteine residues, 1 μL of 350 mM iodoacetamide was added, and the sample was incubated at room temperature in the dark for 30 min. The trypsin digestion reaction was initiated by adding 60 μL of 50 mM Tris-HCl, pH 8.0 and 1 μL of 100 ng μL^{-1} sequencing-grade trypsin (Promega, cat. no. V511). The sample was incubated at 37 $^\circ\text{C}$ for 16 h, immediately desalted by Millipore C₁₈ ZipTip, and analyzed by MALDI mass spectrometry. Data were acquired on a Bruker UltrafleXtreme MALDI-TOF mass spectrometer with matrix 2,5-dihydroxybenzoic acid in positive ion mode. Peptide fragments were calculated using ExPASy. No phosphorylation products peaks were observed for all identifiable Tyr-containing peptide fragments (Fig. S3, ESI†).

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